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(54) THE: RECOMBINANT LIPAGE AND ALPHA-AMYLAGE VARIANTS

(57) Abstract

The present invention relates to lipuse and o-empires various, stabilized towards the inactivation caused by precisions system, in which various a astumily occasing tyrosine neither has been deleted or substituted with a different amino acid residue at one or more which various a astumily occasing tyrosine neither has been deleted or substituted with a different amino acid residue at the inactivation caused by precisions positions.

The invention also relates to a method of stabilizing a lipuse and/or an o-empire various of the invention.

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RECONBINANT LIPASE AND ALPHA-ANYLASE VARIANTS

TECHNICAL PIELD

The present invention relates to lipase and amylase variants, stabilized towards the inactivation caused by peroxidase systems, in which lipase and amylase variants a naturally occurring tyrosine residue has been deleted or substituted with a different amino acid residue at one or more positions.

The invention also relates to a method of stabilis-10 ing a lipase or an α -amylase towards the inactivation caused by peroxidase systems, and detergent compositions comprising a lipase and/or an α -amylase variant of the invention.

RACKGROUND ART

Peroxidases (E.C. 1.11.1.7) are enzymes that catalyse the oxidation of a substrate (an electron or hydrogen donor) with hydrogen peroxide. Such enzymes are known from microbial, plant and animal origins, e.g. peroxidase from Coprinus cinereus (cf. e.g. RP Patent Application 179,486). They are typically hemoproteins, i.e. they contain a heme as a prosthetic group.

Use of peroxidase together with hydrogen peroxide or a hydrogen peroxide precursor has been suggested e.g. in bleaching of pulp for paper production, in treatment of waste water from pulp production, for improved bleaching in laundry s detergents, for dye transfer inhibition during laundering, and for lignin modification, e.g. in particle board production.

Peroxidase systems (also designated POD systems)
comprising a peroxidase or a compound exhibiting peroxidase
activity, a source of hydrogen peroxide, and a peroxidase
menhancing agent, are used for preventing coloured substances,
which leach from dyed fabrics, to deposit on other fabrics
present in the same wash (this phenomenon is commonly known as

dya transfer). Detergent compositions or wash liquors comprising such peroxidase systems have been described in e.g. International Patent Applications WO 92/18687 and WO 92/18683.

A major drawback in applying such peroxidase systems to detergent compositions is that the enzymes present in such compositions may be strongly affected by the peroxidase system, thereby hampering the washing performance of the detergent composition.

SUICKARY OF THE INVENTION

It has now surprisingly been found that lipases and α -amylases may be stabilized towards inactivation caused by peroxidase systems, by deletion or substitution of one or more naturally occurring tyrosine residues with a different amino acid residue.

Accordingly, the invention provides a lipase and/or an α -amylase variant, in which one or more naturally occurring tyrosine residues have been deleted or substituted with a different amino acid residue.

In another aspect, the invention provides a mathod m of stabilization of a lipase and/or an a-amylase variant towards inactivation caused by a peroxidase system, in which method one or more naturally occurring tyrosine residues are deleted or substituted with a different amino acid residue.

In a further aspect, the invention provides deters gent compositions comprising a lipase and/or an a-amylase variant of the invention.

In a yet further aspect, the invention provides detergent additives comprising a lipase and/or an α -amylase variant of the invention.

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DETAILED DISCLOSURE OF THE INVENTION

The present invention provides novel lipase and q-amylase variants, stabilized towards inactivation caused by peroxidase systems.

In the context of this invention, a stabilized lipase or α -amylase variant is a lipase or an α -amylase having improved stability towards inactivation caused by peroxidase systems, when compared to the parent lipase or α -amylase.

Amino Acids

As abbreviations for amino acids the following symbols are used:

~				,	
	λ	-	λla	· 🛥	Alanine
	C	=	Cys	===	Cysteine
	D	**	Asp	=	Aspartic acid
	E	-	Glu		Glutamic acid
	F	· 🗪	Phe	.	Phenylalanine
	G	***	Gly	D16	Glycine
	H		His		Histidine
	Ţ	-	Ile	-	Isoleucine
	K	. ===	Lys	-	Lysine
	I,	=	Leu	=	Leucine
	X	. 20	Met		Methionine
	N	862	Asn		Asparagine
	P	-	Pro		Proline
	Q	=	Gln	-	Glutamine
	R		Arg	•	Arginine
٠.	8	, 488	Ser	-	Serine
	T	-	Thr		Threonine
	V	-	Val	and a	Valine
	W	. =	Trp		Tryptophan
	Y	-	Tyr	108	Tyrosine
:	••	•		•	
•	B	· 🖚	Aex	53 '•	Asp (D) or Asn (N)
	Z	. =	Glx	_	Glu (E) or Gln (Q)
	*				are (2) or arm (6)
	x	=	an arbi	trary a	mino acid
	•		deleti	arl a	sent amino acid
			المراحة المراج	UL GD	GENT ANTHO ACTO :

Peroxidase Activity

In the context of this invention, the enzymatic activity of peroxidases is expressed in "Peroxidase Units" (PODU). In the presence of hydrogen peroxide peroxidases (E.C.

1.11.1.7) catalyse the dehydrogenation of 2,2'-azino-bis (3-ethylbenzothiazoline-6-sulfonate) (ABTS). The greenish-blue colour produced is monitored photometrically at 418 mm. One PODU is the amount of enzyme which, under standard conditions (i.e. pH 7.0; hydrogen peroxide as substrate; 0.1 M phosphate buffer; an incubation temp. of 30°C; an incubation time of 3 min. measured kinetically) catalyses the conversion of 1 mmol of hydrogen peroxide per minute.

Lipase Activity

In the context of this invention, the enzymatic activity of lipases is expressed in Lipase Units. A Lipase Unit (LU) is the amount of enzyme which under standard conditions, i.e. 30.0°C; pH 7.0; tributyrine substrate, liberates 1 µmol titratable butyric acid per minute.

15 g-amylase Activity

The α-amylase activity is measured as absorption/ml at 620 nm using Phadebas tablets (Phadebasv Amylase Test; Pharmacia Diagnostics, SW). The assay is carried out at 60°C.

Peroxidase Systems

In the context of this invention, a peroxidase system is a system comprising a peroxidase or a compound exhibiting peroxidase activity, a source of hydrogen peroxide, and a peroxidase enhancing agent. Such peroxidase systems have been used for obtaining a dye transfer inhibition and have been described in e.g. International Patent Applications WO 92/18687 and WO 92/18683.

In such a peroxidase system, the peroxidase or the compound exhibiting peroxidase activity may be any peroxidase comprised by the enzyme classification EC 1.11.1.7, or any fragment derived therefrom, exhibiting peroxidase activity, or synthetic or semisynthetic derivatives thereof (e.g. porphyrin ring systems or microperoxidases, cf. e.g. US Patent 4,077,768, EP Patent Application 537,381, International Patent Ap-

plications 80 91/05858 and 80 92/16634). Such paroxidades are known from Discrebial, plant and animal origins.

Moreoradian or soy bean percuidase) or alcroorganisms such as fungi or sactoria. Some prediction or alcroorganisms such as fungi or sactoria. Some prediction fungi include strains belonging to the subdivision function, class symbosycotiae, e.g. function, subdivision function, whereas, o.g. function, subdivision function, subjection, function, arthurwises, calentions, subjection, function, subdivision function, subjection, calention function or function, in particular function function, substituted insolars, substituted function, substituted fun

other preferred fungi include strains belonging to the subdivision Basidicaycotina, class Basidicaycotes, e.g. Convinus, Phanexochaete, Coniolus or Transtas, in particular Convinus cinarums for microspanus (IFO 8371), Convinus pagrore bigus, Phanoxochaete chrysospanium (e.g. NA-12) or Transtas, a (previously called Folypanus), e.g. Townsianlar (e.g. FRA 28-A).

Purther preferred fungi include strains balonging to the subdivision zygosycotina, class zycoraceae, e.g. Rhironna or Mucor, in particular Mucor bienalis.

Some preferred bacteria include strains of the order Actinomycotales, e.g. <u>Streptonycos soheroides</u> (AFTC 23963), <u>Streptonycos ibermoviolacens</u> (XFO 12382) or <u>Streptonycos ibermoviolacens</u> (XFO 12382) or <u>Streptonycos ibermoviolacens</u>

Other preferred bacteria include <u>Baciling mailing</u>

12 (ATCC 12905), <u>Bacilius sterrothermonhilus</u>, <u>Rhodobacter acheerpides, Rhodomonas palustri, <u>Streptococcus lactis, Pacudomonas</u>

12 purrecinia (ATCC 15958) or <u>Pseudomonas Rluoroscena</u> (NFEL 8-11)

13 Further proferred bacteria include strains balonging

14 to Myrococcus, e.g. <u>M. virescens</u></u>

Other potential sources of useful particular paronidases are listed in <u>Saunders B C</u>, <u>op, clt.</u>, pp. 41-43.

The percentions may furthermore be one which is producible by a method comprising cultivating a bost call transformed with a recombinant DNA vector which carries a DNA sequence encoding caid percentions as well as DNA sequences ancoding the perceitaing the empression of the DNA sequence encoding the perceitase, in a culture media which conditions peralting the expression of the perceidance and recovering the perceidance from the culture.

Particularly, a recombinantly produced peroxidance is no a peroxidance derived from a <u>Coprimus</u> sp., in particular <u>C.</u> <u>Bacrophizus</u> or <u>C. cineraus</u> according to WO 92/16634.

In the context of this invention, compounds or hibiting paroxidance activity comprise percuidance active fragments derived from cytochromes, becoglobin or percuidance analysis, and synthetic or semisynthetic derivatives thereof, e.g. iron perphins, iron perphyrins, and iron phthalogyanine and derivatives thereof.

In a peroxidate system, the enhancer may be an oxidizable substrate e.g. metal ions or phonolic especiand couch as 7-hydroxycowarin (780m), vanillin (VAN), and p-hydroxycowarincateness (pensyl operational in c.g. International Patent Applications NO 92/18683 and NO 92/18687, and Kate M and Shirixy S, Plant Cell Physiol. 1985 26 (7), pp. 1291-1301 (effections in particular), and Savaders B C, et al., Forestidated In London, 1964, p. 141 ff. or 2,21-asino-bis (3-ethylbanzo-thiazoline-6-sulfonate) (AETS), described in applicant of copending OK Patent Application No. 9201441.

MIGREGE

In a proferred embediment, the lipace of the minution is obtainable from a strain of Empirela, e.g. E. Lamusinosa, E. braviscora, E. bravis var. thermoides, or E. insolens. Lipases obtainable from Empirela are described in e.g. Us Patent 4,810,414, EP Application 305,216 and International Patent Application WO 89/01969, which publications examinably included by reference.

In another specific embodiment, the lipase is obtainable from a strain of <u>Pseudomonas</u>, e.g. <u>Ps. cepacia</u>; <u>Ps. fragi</u>, <u>Ps. stutzeri</u>, or <u>Ps. fluorescens</u>. Lipases obtainable from <u>Pseudomonas</u> are described in e.g. International Patent Publication 89/04361, which publication is hereby included by reference.

In a third specific embodiment, the lipase is obtainable from a strain of <u>Fusarium</u>, e.g. <u>F. oxysporum</u>. Lipases obtainable from <u>Fusarium</u> are described in e.g. EP Publication 130,064 and EP Publication 395,678, which publications are hereby included by reference.

In further specific embodiments, the lipase is obtainable from a strain of Rhizomucor, s.g. Rhizomucor mishei, or a strain of Candida, e.g. C. antarctica, or C. cylindraces s (also called C. rugosa), or a strain of Chromobacterium, e.g. C. viscosum.

In a more preferred embodiment, a lipase variant of the invention is a <u>Humicola lammginosa</u> lipase having an amino acid sequence as described in EP Publication 305,216 (in which me publication the amino acid sequence is presented in Fig. 5), which sequence has been changed in one or more of the following positions: 16, 21, 53, 138, 164, 171, 194, 213, 220, 261.

Amylases

In a preferred embodiment, the a-amylase variant of state invention is obtainable from a strain of <u>Bacillus</u> or a strain of <u>Aspervillus</u>.

In a more specific embodiment, the α -amylase variant is obtainable from a strain of <u>B. licheniformis</u>. The amino acid sequence for the <u>B. licheniformis</u> 584 α -amylase (<u>Stephens et mal.</u>) appears from J. Bacteriol. 1984 158 369-372, and J. Bacteriol. 166, 635-643, 1986, FR 2665178 or EP 410498. Thus, the tyrosine positions are: 10, 14, 31, 46, 56, 59, 62, 77, 98, 150, 158, 175, 193, 195, 198, 203, 219, 262, 273, 290, 302, 348, 358, 363, 367, 394, 396, 402, 439, 480.

In another specific enbodiment, the e-orginal variant is obtainable from a strain of <u>B. anyloliquefactens</u>. The orino acid sequence for the <u>B. anyloliquefactens</u> a-orginal (Takkinen at al.) appears from J. Biel Chem. 1983 258 1007-1013.

In a third specific embeditiont, the e-maylase variant is obtainable from a strain of <u>R. stranothermontiles</u>. The amino acid sequence for the <u>R. stranothermontiles</u> e-maylase appears from J. Bacteriol. 166, 635-643, 1986.

In a fourth specific embediesnt, the e-aryland wariant is obtainable from a strain of <u>A. miger</u>. Was aring cold sequence for the <u>A. miger</u> e-aryland appears from DK Fotomt Application 5126/87.

In further specific embodinents, e-anylase variants of the invention are chimeric e-anylases. Chimeric e-anylases s are disclosed in e.g. EP Potent Publication 252,666.

Hethods of Stabilizing Lipases and a-paylases

The present invention provides a nethed of stabilizing lipuses and s-applaces towards inactivation cancel by
peroxidase systems, by which method one or nore naturally
a occurring tyrosine residues are deleted or substituted with a
different amino acid residue.

Recombinantly Produced Manses and c-amplanes

In the past, numerous processes have been developed for the production of polypoptides or proteins by mans of the recombinant DMA technology. Hostly used for this purpose are E. coli. Encillus subtilis, Enccinrovers cerevising and different Aspervillus strains, e.g. A. organ and A. nigg. Especially the Aspervillis are attractive candidates as host microorganisms for recombinant DMA vectors being vall-characterized and videly used microorganisms for the commercial production of enzymes. In Aspervillus organism, estheds have been developed for transformation of the organism, and production of several enzymes, among these the Numicola lanualness and Phisometer michai lipases (vide e.g. European Patent Applications 238,023

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and 305,216, and International Fatant Application We 89/01969), which publications are bearby included by reformes.

Thoresion of Polyneptides Blosynthetically

Upon transformation of an organism where the intenition is production of a polypeptide or a protein, a DNA
sequence is introduced into the organism. The sequence contains
the coding region of the gene of interest flanked by transcription/translation start signals and transcription/translation
termination signals. The coding region contains units of three
house pairs, called codons; which upon translation of the
transcribed gene are translated into asino acids, which again
are assembled to give the polyperation of interest.

Introducing Mutations in Polymortidas

By changing one or Bore specific ecdons in the coling 15 region and transforming the host microorganism with these ner coding regions, now polypaytides can be protocod which differ from the original polypeptide by one or more anino acids. Such alterations can be introduced by beams of a technique generally known as "site-directed in vitro entagenesis". A number of w methods have been published. An early method is described by <u>zollar & Szith,</u> DNA 1984 I (6) 479-488, azd izvolvec use of the singlo-stranded KD3 bacteriophago. A preforred nethed using FC: (polyreruse chain reaction) is described by Nelson (News) Analytical Biochemistry, 1989 100 147-151. It involves a 1-stop by denotation of a PCR fragment containing the objection by valiag a chemically synthesized DNA oligonvalectics as one of the primary in the FCR reactions. From the FCR-generated fragment, a DNA fragment carrying the mutation can be incluted by cleavage with restriction enzymes and re-inserted into the w expression plassid. A third nutagenesis method takes advantage of restriction sites in the DNA coding region. By digasting the DAN vith restriction enzymes at mitee flanking the mutagenesis target, synthesizing a nev fragment synthetically containing the desired mutation and cloning this new fragment between the restriction sites, a mutant coding region can be constructed.

All methods are generally applicable to investigations in the field called protein engineering which deals with the development of polypeptides with new or altered characteristics.

Transformation and expression may be accomplished by methods known in the art, e.g. as described in European Patent Application 305,216, which specification is hereby included by reference.

The microorganisms able to produce a stabilized lipase or a-amylase of this invention can be cultivated by conventional fermentation methods in a nutrient medium containing assimilable carbon and nitrogen together with other sessential nutrients, the medium being composed in accordance with the principles of the known art. Purification and recovery of the stabilized lipase or a-amylase may also be conducted in accordance with methods known per se.

Nucleotide Sequences, Expression Vectors And Microorganisms

This invention also relates to DNA nucleotide sequences encoding a stabilized lipase or α-amylase of the invention. The stabilized lipase or α-amylase may be expressed and produced when DNA nucleotide sequence encoding the lipase or α-amylase is inserted into a suitable vector in a suitable shost organism. The host organism is not necessarily identical to the organism from which the parent gene originated. The construction of the mutated genes, vectors and mutant and transformed microorganisms may be carried out by any appropriate recombinant DNA technique, known in the art.

The invention also relates to expression vectors and host organisms containing a DNA nucleotide encoding a stabilized lipase or α -amylase of this invention.

Determent Compasitions

According to the invention, the lipace and the applane variant may typically be a component of a detergent composition. As such, it may be included in the determent s composition in the form of a non-dusting granulate, a stabilised liquid, or a protected enzyee. Non-dusting granulates usly be produced, a.g., as disclosed in US 4,106,991 and 4,661,452 (both to nove industri L/S) and may optionally be coated by matheds known in the art. Examples of rang coating materials (per poly (ethylene oxide) products (polyethyleneglyce), Fet with mean molar weights of 1000 to 20000, ethoxylated nonylphonols having from 16 to 50 ethylene oride units; ethorylated fatty alcohols in which the alcohol contains from 12 to 30 carbon atoms and in which there are 15 to 80 ethylene exide 15 units: fatty alcohols: fatty acids: and mono- and di- and triglycarides of fatty acids. Bramples of film-forming coating materials suitable for application by fluid bed techniques are givon in patent CB 1483591. Liquid enzyme preparations may, for instance, be stabilized by adding a polyol such as propylers m glycol, a sugar or sugar alcohol, lactic acid or boric acid according to established pethods. Other enzyme stabilizers are woll known in the art. Protected enzyres may be prepared according to the method disclosed in MP 238, 216.

The detergent composition of the invention may be in any convenient form, e.g. as porder, granulas, pasts or liquid. A liquid detergent may be aqueous, typically contains up to 70 % rater and 0-30 % organic solvent, or nonagueous.

The detergent composition comprises one or note surfactants, each of which may be anionic, nonionic, cationic, or switterionic. The detergent will usually contain 0-50 % of anionic surfactant such as linear allybenzenesulfonate (LAS), alpha-olefinsulfonate (AOS), alkyl sulfate (fatty alcohol sulfate) (AS), alcohol ethorysulfate (AEOS or AES), secondary alleaseulfonates (SAS), alpha-sulfo fatty acid methyl esters, allyl- or alkenylsuccinic acid or soap. It may also contain de-

40 % of nonionic surfactant such as alcohol ethoxylate (ABO or AB), carboxylated alcohol ethoxylates, nonylphenol ethoxylate, alkylpolyglycoside, alkyldimethylamineoxide, ethoxylated fatty acid monoethanolamide, fatty acid monoethanolamide, alkyl-(N-5 methyl)-glucoseamide or polyhydroxy alkyl fatty acid amide (e.g. as described in WO 92/06154).

The detergent composition may additionally comprise one or more other enzymes, such as cutinase, protease, cellulase, peroxidase, or oxidase.

The detergent may contain 1-65 % of a detergent builder or complexing agent such as reolite, diphosphate, triphosphate, phosphonate, citrate, nitrilotriacetic acid (NTA), ethylenediaminetetraacetic acid (EDTA), diethylenetriaminepentaacetic acid (DTMPA), alkyl- or alkenylsuccinic acid, soluble silicates or layered silicates (e.g. SKS-6 from Hoechst). The detergent may also be unbuilt, i.e. essentially free of detergent builder.

The detergent may comprise one or more polymers.

Examples are carboxymethylcallulose (CMC), poly(vinylm pyrrolidone) (PVP), polyethyleneglycol (PEG), poly(vinylalcohol) (PVA), polycarboxylates such as polyacrylates,
maleic/acrylic acid copolymers and lauryl methacrylate/acrylic
acid copolymers.

The detergent may contain a bleaching system which may comprise a H₂O₂ source such as perborate or percarbonate which may be combined with a peracid-forming bleach activator such as tetraacetylethylenediamine (TAKD) or nonanoyloxyben-zenesulfonate (NOBS). Alternatively, the bleaching system may comprise peroxyacids of e.g. the amide, imide, or sulfone with the system and the system and the system peroxyacids of e.g. the amide, imide, or sulfone type.

The enzymes of the detergent composition of the invention may be stabilized using conventional stabilizing agents, e.g. a polyol such as propylene glycol or glycerol, a sugar or sugar alcohol, lactic acid, boric acid, or a boric acid derivative as e.g. an aromatic borate ester, and the com-

position may be formulated as described in e.g. WO 92/19709 and WO 92/19708.

The detergent may also contain other conventional detergent ingredients such as e.g. fabric conditioners inscluding clays, foam boosters, suds suppressors, anti-corresion agents, soil-suspending agents, anti-soil redeposition agents, dyes, bactericides, optical brighteners, or perfume.

The pH (measured in aqueous solution at use concentration) will usually be neutral or alkaline, e.g. 7-11.

Particular forms of detergent compositions within the scope of the invention include:

1) A detergent composition formulated as a granulate having a bulk density of at least 600 g/l comprising

- linear alkylbenzenesulfonate (calculated as acid)	7 - 12%
- alcohol ethoxysulfate (e.g. C ₁₂₋₁₆ alcohol, 1-2 EO) or alkyl sulfate (e.g. C ₁₄₋₁₈)	1 - 48
- alcohol ethoxylata zo (e.g. C ₁₄₋₁₅ alcohol, 7 EO)	5 - 9%
- sodium carbonate (as Ma ₂ CO ₃)	14 - 20%
- soluble silicate (as Na ₂ O,2SiO ₂)	2 - 6%
- zeolite (as NaAlSiO ₄)	15 - 224
- sodium sulfate (as Na ₂ SO ₄)	0 - 6%
 sodium citrate/citric acid (as C,H,Na,O,/C,H,O,) sodium perborate (as NaBO, H,O) 	0 - 15 t 11 - 18 t
- TAED	3 - 6\$
- carboxymethylcellulose	0 - 2%
m - polymers (e.g. maleic/acrylic acid copolymer, PVP, PEG)	0 - 3%
- enzymes	0 - 5%
- minor ingredients (e.g. suds supressors, perfume, optical brightener, photobleach)	
a set Americe (Princented CII)	0 - 5%

2) A detergent composition formulated as	a granulate having a
bulk density of at least 600 g/l comprisi	
 linear alkylbenzenesulfonate (calculated as acid) 	6 - 11\$
s - alcohol ethoxysulfate (e.g. C ₁₂₋₁₈ alcohol, 1-2 E0) or alkyl sulfate (e.g. C ₁₄₋₁₈)	1 - 38
 alcohol ethoxylate (e.g. C_{N-15} alcohol, 7 EO) 	5 - 9%
10 - sodium carbonate (as Na ₂ CO ₃)	15 - 21%
- soluble silicate (as Na ₂ 0,2SiO ₂)	1 - 48
- zeolite (as NaAlSiO ₄)	24 - 34%
- sodium sulfate (as Na ₂ SO ₄)	4 - 10%
- sodium citrate/citric acid is (as C ₆ H ₅ Na ₅ O ₇ /C ₆ H ₆ O ₇) - carboxymethylcellulose	0 - 15 % 0 - 2 %
- polymers (e.g. maleic/acrylic acid cop PVP, PEG)	olymer, 1 - 6%
- enzymes	0 - 5%
<pre>a - minor ingredients (e.g. suds supressors, perfume)</pre>	0 - 5%
3) A detergent composition formulated a bulk density of at least 600 g/l compris	
- linear alkylbenzenesulfonate calculated as acid)	5 - 9%
<pre>- alcohol ethoxylate (e.g. C₁₂₋₁₅ alcohol, 7 EO)</pre>	7 - 14%
- soap as fatty acid (e.g. C ₁₄₋₂₂)	1 - 3%
30 - sodium carbonate (as Na ₂ CO ₃)	10 - 178
- soluble silicate (as Na ₂ 0,2SiO ₂)	3 - 9%
- zeolite (as NaAlSiO _L)	23 - 33%
- sodium sulfate (as Na ₂ SO ₄)	0 - 4%

- sodium perborate (as NaBO3.H2O)	8 - 16%
- TAED	2 - 8%
- phosphonate (e.g. EDTMPA)	0 - 1%
- carboxymethylcallulose	0 - 2%
 polymers (e.g. maleic/acrylic acid copolymers, PEG) 	1 - 3%
- enzymes	0 - 5%
 minor ingredients (e.g. suds supressors, perfume, optical brightener) 	0 - 5%
04) A detergent composition formulated as a bulk density of at least 600 g/l comprising	
 linear alkylbenzenesulfonate (calculated as acid) 	8 - 12%
- alcohol ethoxylate s (a.g. C ₁₂₋₁₃ alcohol, 7 EO)	10 - 25%
- sodium carbonate (as Na ₂ CO ₃)	14 - 228
- soluble silicate (as Na ₂ O, 2SiO ₂)	1 - 5%
- zeolite (as NaAlSiO4)	25 - 35%
- sodium sulfate (as Na ₂ SO ₄)	0 - 10%
c - carboxymethylcellulose	0 - 2%
 polymers (e.g. maleic/acrylic acid copolyr PVP, PEG) 	mer, 1 - 3%
- enzymes	o – 5%
- minor ingredients (e.g. suds supressors, perfume)	0 - 5%
5) An aqueous liquid detergent composition	comprising
 linear alkylbenzenesulfonate (calculated as acid) 	15 - 21%
- alcohol ethoxylate (e.g. C ₁₂₋₁₅ alcohol, 7 EO or C ₁₂₋₁₅ alcohol, 5 EO)	12 - 18%

- soap as fatty acid (e.g. oleic acid)	3 - 138
- alkenylsuccinic acid (C12-14)	0 - 13%
- aminoethanol	8 - 18%
- citric acid	2 - 8%
5 - phosphonate	0 - 3%
- polymers (e.g. PVP, PEG)	0 - 3%
- borate (as B ₄ O ₇)	0 - 2%
- ethanol	0 - 3%
- propylene glycol	8 - 14%
10 - enzymes	0 - 5%
 minor ingredients (e.g. dispersants, suds supressors, perfume, optical brightener) 	0 - 5%
6) An aqueous structured liquid detergen	composition compris
 6) An aqueous structured liquid detergent ing linear alkylbenzenesulfonate 	
 6) An aqueous structured liquid detergents ing linear alkylbenzenesulfonate (calculated as acid) alcohol ethoxylate (e.g. C_{12.55} alcohol, 7 EO 	15 - 218
 6) An aqueous structured liquid detergents ing linear alkylbenzenesulfonate (calculated as acid) alcohol ethoxylate (e.g. C₁₂₋₁₅ alcohol, 7 EO or C₁₂₋₁₅ alcohol, 5 EO) 	15 - 21 % 3 - 9 %
 6) An aqueous structured liquid detergents ing linear alkylbenzenesulfonate (calculated as acid) alcohol ethoxylate (e.g. C₁₂₋₁₅ alcohol, 7 EO or C₁₂₋₁₅ alcohol, 5 EO) soap as fatty acid (e.g. oleic acid) 	15 - 21% 3 - 9% 3 - 10%
 6) An aqueous structured liquid detergents ing linear alkylbenzenesulfonate (calculated as acid) alcohol ethoxylate (e.g. C₁₂₋₁₅ alcohol, 7 EO or C₁₂₋₁₅ alcohol, 5 EO) soap as fatty acid (e.g. oleic acid) zeolite (as NaAlSiO₄) 	15 - 21% 3 - 9% 3 - 10% 14 - 22%
 6) An aqueous structured liquid detergents ing linear alkylbenzenesulfonate (calculated as acid) alcohol ethoxylate (e.g. C₁₂₋₁₅ alcohol, 7 EO or C₁₂₋₁₅ alcohol, 5 EO) soap as fatty acid (e.g. oleic acid) zeolite (as NaAlSiO₄) potassium citrate borate (as B₄O₇) 	15 - 21% 3 - 9% 3 - 10% 14 - 22% 9 - 18%
 6) An aqueous structured liquid detergents ing linear alkylbenzenesulfonate (calculated as acid) alcohol ethoxylate (e.g. C₁₂₋₁₅ alcohol, 7 EO or C₁₂₋₁₅ alcohol, 5 EO) soap as fatty acid (e.g. oleic acid) zeolite (as NaAlSiO₄) potassium citrate borate (as B₄O₇) 	15 - 21% 3 - 9% 3 - 10% 14 - 22% 9 - 18% 0 - 2%
6) An aqueous structured liquid detergents ing - linear alkylbenzenesulfonate (calculated as acid) - alcohol ethoxylate (e.g. C ₁₂₋₁₅ alcohol, 7 EO or C ₁₂₋₁₅ alcohol, 5 EO) - soap as fatty acid (e.g. oleic acid) - zeolite (as NaAlSiO ₄) - potassium citrate - borate (as B ₄ O ₇) z - carboxymethylcellulose	15 - 21% 3 - 9% 3 - 10% 14 - 22% 9 - 18% 0 - 2% 0 - 2% 0 - 3%
 6) An aqueous structured liquid detergents ing linear alkylbenzenesulfonate (calculated as acid) alcohol ethoxylate (e.g. C₁₂₋₁₅ alcohol, 7 EO or C₁₂₋₁₅ alcohol, 5 EO) soap as fatty acid (e.g. oleic acid) zeolite (as NaAlSiO₄) potassium citrate borate (as B₄O₇) carboxymethylcellulose polymers (e.g PEG, PVP) anchoring polymers as e.g. lauryl metharylate/acrylic acid of 	15 - 21% 3 - 9% 3 - 10% 14 - 22% 9 - 18% 0 - 2% 0 - 2% 0 - 3%

bulk density of at least 600 g/l comprising - fatty alcohol sulfate - ethoxylated fatty acid monoethanolamide - soap as fatty acid - sodium carbonate (as Na ₂ CO ₃) - soluble silicate (as Na ₂ O ₃ CSiO ₂) - zeolite (as NaALSiO ₄) - sodium sulfate (as Na ₂ SO ₄) - sodium perborate (as NaBO ₃ .H ₂ O) - TAED - polymers (e.g. maleic/acrylic acid copol PEG) - enzymes - minor ingredients (e.g. optical brighten suds supressors, perfume) 8) A detergent composition formulated as a clinear alkylbenzenesulfonate (calculated as acid) - ethoxylated fatty acid monoethanolamide - soap as fatty acid - sodium carbonate (as Na ₂ CO ₃) - soluble silicate (as Na ₂ CO ₃) - soluble silicate (as Na ₂ SO ₄) - sodium sulfate (as Na ₂ SO ₄) - sodium citrate (as C ₆ H ₅ Na ₃ O ₇)	ume, 0 - 5%			
7) A detergent composition formulated as a	-			
-				
- fatty alcohol sulfate	5 - 10%			
- ethoxylated fatty acid monoethanolamida	3 - 9%			
- soap as fatty acid	0 - 3%			
- sodium carbonate (as Na ₂ CO ₃)	5 - 10%			
10 - soluble silicate (as Na ₂ O,2SiO ₂)	1 - 4%			
- zeolite (as NaAlSiO ₄)	20 - 40%			
- sodium sulfate (as Na _z SO ₄)	2 - 84 .			
- sodium perborate (as NaBo ₃ .H ₂ O)	12 - 18%			
- TAED	2 - 7%			
s - polymers (e.g. maleic/acrylic acid copoly PEG)	mer, 1 - 5%			
- enzymes	0 - 5%			
- minor ingredients (e.g. optical brightene suds supressors, perfume)	r, 0 - 5%			
zo 8) A detergent composition formulated as a gr	ranulate comprisi			
	8 - 14%			
- ethoxylated fatty acid monoethanolamide	5 - 11%			
- soap as fatty acid	0 - 3%			
s - sodium carbonate (as Na ₂ CO ₃)	4 - 10%			
- soluble silicate (as Na ₂ O,2SiO ₂)	1 - 4%			
- zeolite (as NaAlSiO4)	30 - 50%			
- sodium sulfate (as Na ₂ SO ₄)	3 - 11%			
- sodium citrate (as C ₆ H ₅ Na ₅ O ₇)	. 5 - 12%			
- polymers (e.g. PVP, maleic/acrylic acid copolymer, PEG)	1 - 5%			

- enzyaes	0 - 5%
 minor ingredients (e.g. suds supressors, perfune) 	0 - 5%
9) A detergent composition formulated as a gr	anulate comprisi
5 - linear alkylbenzenesulfonate (calculated as acid)	6 - 12\$
- nonionic surfactant,	1 - 48
- soap as fatty acid	2 - 6%
- sodium carbonate (as Na ₂ CO ₃)	14 - 228
10 - zeolite (as NaAlSiO4)	18 - 32\$
- sodium sulfate (as Na _Z SO ₄)	5 - 20%
- sodium citrate (as C ₆ H ₅ Na ₃ O ₇)	3 - 8\$
- sodium perborate (as NaBO3.H2O)	4 - 98
- bleach activator (e.g. NOBS or TAED)	1 - 5%
ts - carboxymethylcellulose	0 - 2%
- polymers (e.g. polycarboxylate or PEG)	1 - 5%
- enzymes	0 - 5%
- minor ingredients (e.g. optical brightener, perfume)	0 - 5%
zo 10) An aqueous liquid detergent composition	n comprising
- linear alkylbenzenesulfonate (calculated as acid)	15 - 23%
- alcohol ethoxysulfate (e.g. C ₁₂₋₁₅ alcohol, 2-3 EO)	8 - 15 t
z - alcohol ethoxylate (e.g. C ₁₂₋₁₅ alcohol, 7 EO or C ₁₂₋₁₅ alcohol, 5 EO)	3 - 9%
- soap as fatty acid (e.g. lauric acid)	0 - 3%
- aminoethanol	1 - 54
w - sodium citrate	5 - 10%

- hydrotrope (e.g. sodium toluenesulfonate)	2 - 6\$	•
- borate (as B ₄ O ₇)	0 - 2%	
- carboxymethylcellulose	0 - 1%	
- ethanol	1 - 38	
s - propylene glycol	2 - 5%	
- enzymes	0 - 5%	
 minor ingredients (e.g. polymers, dispersar perfume, optical brighteners) 	ots, 0 - 5 t	٠.
11) An aqueous liquid detergent composition of	comprising	
s - linear alkylbenzenesulfonate (calculated as acid)	20 - 32%	
- alcohol ethoxylate (e.g. C ₁₂₋₁₅ alcohol, 7 EO or C ₁₂₋₁₅ alcohol, 5 EO)	6 - 12 %	
5 - aminosthanol	2 - 6%	
- citric acid	8 ~ 14%	
- borate (as B ₄ O ₇)	1 - 38	
 polymer (e.g. maleic/acrylic acid copolymer anchoring polymers as e.g. lauryl methacrylate/acrylic acid copolymer and CMC) 	•	
- glycerol	0 - 3 1	
- enzymes	0 - 5%	,
- minor ingredients (e.g. hydrotropes, dispersants, perfume, optical brighteners)		
12) A detergent composition formulated as a gr		ge
bulk density of at least 600 g/l comprisi	ng	
 anionic surfactant (linear alkylbenzenesulfonate, alkyl sulfate, alpha olefinsulfonate, alpha-sulfo fatty acid methyl esters, alkanesulfonates, soap) 	- 25 - 40 \	
- nonionic surfactant (e.g. alcohol ethorylate)		
re.g. alcondi ethoxviata)	1 - 102	

- sodium carbonate (as Na ₂ ∞ ₃)	8 - 25%
- soluble silicates (as Na ₂ O, 2SiO ₂)	5 - 15%
- sodium sulfate (as Na _z SO ₄)	0 - 5%
- zeolita (as NaAlSiO4)	15 - 28\$
5 - sodium perborate (as NaBO ₃ .4H ₂ O)	0 - 20%
- bleach activator (TAED or HOBS)	0 - 54
- enzymes	0 - 5%
 minor ingredients (e.g. perfume, optical brighteners) 	0 - 3\$

- 10 13) Detergent formulations as described in 1) 12) where the content of linear alkylbenzenesulfonate or a part of it is substituted by alkyl sulfate $(C_{12}-C_{18})$.
- 14) Detergent formulations as described in 1) 13) which contain a stabilized or encapsulated peracid either as an additional component or as a substitute for already specified bleach systems.
 - 15) Detergent compositions as described in 3), 7), 9) and 12) where the content of perborate is substituted by percarbonate.
- 16) Detergent composition formulated as a nonaqueous detergent 20 liquid comprising a liquid nonionic surfactant as e.g. linear alkoxylated primary alcohol, a builder system (e.g. phosphate), enzyme and alkali. The detergent may also comprise anionic surfactant and/or a bleach system.

The lipase and c-amylase variants of the invention z may be incorporated in concentrations conventionally employed in detergents. It is at present contemplated that, in the detergent composition of the invention, the lipase and camylase variants may be added in an amount corresponding to 0.001-100 mg of lipase or c-amylase variant per liter of wash

CLAIMS

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- 1. A lipase or an c-amylase variant, stabilized towards inactivation caused by a peroxidase system comprising a peroxidase or a compound exhibiting peroxidase activity, a source of hydrogen peroxide and a peroxidase enhancing agent, characterized in, that one or more naturally occurring tyrosine residues has/have been deleted or substituted with a different amino acid residue.
- 2. A lipase or an α-amylase variant according to 10 claim 1, in which one or more tyrosine residue(s) has/have been substituted with a phenylalanine residue, a leucine residue, an isoleucine residue, a valine residue, a glutamine residue, an asparagine residue, a serine residue, a threonine residue, a glutamic acid residue, or a histidine residue.
 - 3. A lipase variant according to either of claims 1-2, the lipase variant being obtainable from a strain of Humicola, Pseudomonas, Fusarium, Rhizomucor, or Candida.
- 4. A lipase variant according to claim 3, the lipase variant being obtainable from a strain of H. lanuginosa, H. more previsiona, H. brevis var. thermoidea, H. insolens, Ps. cepacia, Ps. fragi, Ps. stutzeri, Ps. fluorescens, P. oxysporum, Rhizomucor miehei, C. antarctica, or C. cylindracea.
 - 5. A lipase variant according to claim 4, the lipase variant being obtainable from a strain of <u>H. lanuginosa</u>.
 - 6. A lipase variant according to claim 5, in which a naturally occurring tyrosine residue has been deleted or substituted in one or more of the following tyrosine positions: 16, 21, 53, 138, 164, 171, 194, 213, 220, 261.

- 7. An α -amylase variant according to either of claims 1-2, the α -amylase variant being obtainable from a strain of Bacillus or Aspergillus.
- 8. An q-amylase variant according to claim 7, the qsamylase variant being obtainable from a strain of B. licheniformis
- 9. An α-amylase variant according to claim 8, in which a naturally occurring tyrosine residue has been deleted or substituted in one or more of the following positions: 10, 14, 31, 46, 56, 59, 62, 77, 98, 150, 158, 175, 193, 195, 198, 203, 219, 262, 273, 290, 302, 348, 358, 363, 367, 394, 396, 402, 439, 480.
- amylase towards inactivation caused by a peroxidase system accomprising a peroxidase or a compound exhibiting peroxidase activity, a source of hydrogen peroxide and a peroxidase enhancing agent, characterized in, that one or more naturally occurring tyrosine residues is/are deleted or substituted with a different amino acid residue.
- by substitution of one or more naturally occurring tyrosine residues with a phenylalanine residue, a leucine residue, an isoleucine residue, a valine residue, a glutamine residue, an asparagine residue, a serine residue, a threonine residue, a glutamic acid residue, or a histidine residue.
 - 12. The method according to either of claims 10-11, the lipase being obtainable from a strain of <u>Humicola</u>, <u>Pseudomonas</u>, <u>Fusarium</u>, <u>Rhizomucor</u>, or <u>Candida</u>.
- 13. The method according to claim 12, the lipase so being obtainable from a strain of <u>H. lanuginosa</u>, <u>H. brevispora</u>,

. .)

H. brevis var. thermoidea, H. insolens, Ps. cepacia, Ps. fraci, Ps. stutzeri, Ps. fluorescens, F. oxysporum, Rhizomucor miehei, C. antarctica, or C. cylindracea.

- 14. The method according to claim 13, the lipase s being obtainable from a strain of <u>Humicola lanuginosa</u>.
 - 15. The method according to claim 14, in which a naturally occurring tyrosine residue has been deleted or substituted in one or more of the following tyrosine positions: 16, 21, 53, 138, 164, 171, 194, 213, 220, 261.
- 16. The method according to either of claims 10-11, the α-amylase being obtainable from a strain of <u>Bacillus</u> or <u>Aspergillus</u>.
 - 17. The method according to claim 16, the q-amylase being obtainable from a strain of B. licheniformis.
- 18. The method according to claim 17, in which a naturally occurring tyrosine residue has been changed in one or more of the following positions: 10, 14, 31, 46, 56, 59, 62, 77, 98, 150, 158, 175, 193, 195, 198, 203, 219, 262, 273, 290, 302, 348, 358, 363, 367, 394, 396, 402, 439, 480.
- 19. A detergent composition comprising a lipage and/or an a-amylase variant according to any of claims 1-9.
- 20. A detergent composition according to claim 19, which further comprises one or more other enzymes, in particular proteases, cellulases, oxidases and/or peroxidases, sometionally used in detergents.
 - 21. A detergent additive comprising a lipase and/or an α -amylase variant according to any of claims 1-9, provided

WO 94/14951 PCT/DK93/00441

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in the form of a non-dusting granulate, a stabilized liquid, a slurry, or a protected enzyme.

International application No. PCT/DK 93/00441

A. CLASSIFICATION OF SUBJECT MATTER		•
IPCS: C12N 9/20, C12N 9/28, C12N 15/55, (According to International Patent Classification (IPC) or to both na	C12N 15/56 // C11D 3/386	
B. FIELDS SEARCHED		
Minimum documentation searched (charification system followed by	chesification symbols)	
IPC5: C12N, C11D Documentation searched other than minimum documentation to the	extent that such documents are included in	the fields marched
•		
SE,DK,FI,NO classes as above		
Historolic data bese consulted during the international search (name	or one one me' was harrene' war	
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WPIL, BIOSIS, CA, EPODOC	f .	
C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category* Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.
P,X WD, A1, 9311254 (NOVO NORDISK A/ (10.06.93), page 5, line 3	/S), 10 Jume 1993 - line 25	1-21
	•	
A EP, A1, 0407225 (UNILEVER PLC), (09.01.91)	9 January 1991	1-21
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A WO, A2, 9100353 (GIST-BROCADES 10 January 1991 (10.01.91)	N.V.),	1-21
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Further documents are listed in the continuation of Bo	ox C. X See patent family and	BSC.
Special categories of check documents: "A" document defining the general state of the set which is set considered.	the principle or theory underlying the	RCHOOF OUR COME IN CONTRACTOR
to be of particular extension "E" refler document but published on or other the international filling date		أرار فوجوشح وجانسو وسيبين
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"P" document published polor to the interestional filling date has later the	period opayons to a bassion serving ar	
the primity date claimed	'&' document number of the same pain. Date of mailing of the international	
Date of the actual completion of the international search	31 -03- 1994	
30 March 1994	01 00 507	
Name and mailing address of the ISA/	Authorized officer	·
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INTERNATIONAL SEARCH REPORT

Information on patent family members

26/02/94 PCT/DK 93/00441

Γ		ocument erch report	Publication date		sober(s)	Publication date	
卜	WO-A1-	9311254	10/06/93	NONE			
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	WO-A2-	9100353	10/01/91	AU-B- AU-A- EP-A-	638263 5953890 0410498	24/06/93 17/01/91 30/01/91	

Porm PCT/ISA/210 (patent family amount) (July 1992)

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